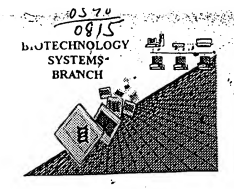
RAW SEQUENCE LISTING ERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number:	19/920262	
Source:	CIPE	
Date Processed by STIC:	08/09/2001	

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,

2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 c-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 c-mail help: patin3help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE <u>CHECKER VERSION 3.0 PROGRAM</u>, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW:

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 – 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST.25.

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2K-compliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money.

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

Raw Sequence Listing Error Summary

ERROR DETECTED	SUGGESTED CORRECTION SERIAL NUMBER: 09/920, 262
ATTN: NEW RULES CASE	S: PLEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE
1 Wrapped Nucleics Wrapped Aminos	The number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it. Please adjust your right margin to .3; this will prevent "wrapping."
2Invalid Line Length	The rules require that a line not exceed 72 characters in length. This includes white spaces.
Misaligned Amino Numbering	The numbering under each 5th amino acid is misaligned. Do not use tab codes between numbers; use space characters, instead.
4Non-ASCII	The submitted file was not saved in ASCII(DOS) text, as required by the Sequence Rules. Please ensure your subsequent submission is saved in ASCII text.
5Variable Length	Sequence(s)contain n's or Xaa's representing more than one residue. Per Sequence Rules, each n or Xaa can only represent a single residue. Please present the maximum number of each residue having variable length and indicate in the <220>-<223> section that some may be missing.
6PatentIn 2.0 "bug"	A "bug" in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid sequences(s) Normally, PatentIn would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. This applies to the mandatory <220>-<223> sections for Artificial or Unknown sequences.
7Skipped Sequences (OLD RULES)	Sequence(s) missing. If intentional, please insert the following lines for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) (i) SEQUENCE CHARACTERISTICS: (Do not insert any subheadings under this heading) (xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) This sequence is intentionally skipped
	Please also adjust the "(ii) NUMBER OF SEQUENCES:" response to include the skipped sequences.
8Skipped Sequences' (NEW RULES)	Sequence(s) missing. If Intentional, please insert the following lines for each skipped sequence. <210> sequence id number <400> sequence id number 000
9Use of n's or Xaa's (NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing. Per 1.823 of Sequence Rules, use of <220>-<223> is MANDATORY if n's or Xaa's are present. In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.
10Invalid <213> Response	Per 1.823 of Sequence Rules, the only valid <213> responses are: Unknown, Artificial Sequence, or scientific name (Genus/species). <220>-<223> section is required when <213> response is Unknown or is Artificial Sequence
11Use of <220>	Sequence(s) missing the <220> "Feature" and associated numeric identifiers and responses. Use of <220> to <223> is MANDATORY if <213> "Organism" response is "Artificial Sequence" or "Unknown." Please explain source of genetic material in <220> to <223> section. (See "Federal Register," 06/01/1998, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of Sequence Rules)
12Patentin 2.0 "bug"	Please do not use "Copy to Disk" function of Patentin version 2.0. This causes a corrupted file, resulting in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing). Instead, please use "File Manager" or any other manual means to copy file to floppy disk.
13Misuse of n	n can only be used to represent a single nucleotide in a nucleic acid sequence. N is not used to represent any value not specifically a nucleotide.

AMC/MH - Biotechnology Systems Branch - 08/21/2001

Does Not Comply Corrected Diskette Needed

DATE: 08/09/2001 TIME: 14:31:25

OIPE

```
Input Set : A:\PTO_VSK.txt
                       Output Set: N:\CRF3\08092001\I920262.raw
       4 <110> APPLICANT: Shealy, David; Knight, David; Scallon, Bernie; Giles-Komar, Jill;
Peritt,
               David
       7 <120> TITLE OF INVENTION: IL-12 ANTIBODIES, COMPOSITIONS, METHODS AND USES
       9 <130> FILE REFERENCE: CEN248
C--> 11 <140> CURRENT APPLICATION NUMBER: US/09/920,262
                                                                      The type of error shown exist throughout
C--> 11 <141> CURRENT FILING DATE: 2001-08-01
                                                                      the Sequence Users and the check subsequent
      11 <160> NUMBER OF SEQ ID NOS (15
      13 <170> SOFTWARE: PatentIn Ver 2-0
                                                                      sequences for the sequences
                                    Errored
                                                                 The type or effors snown exist throughout
ERRORED SEQUENCES
                                                                 the Sequence Listing. Please check subsequent
                                                                 sequences for similar errors.
     15 <210> SEQ ID NO: 1
     16 <211> LENGTH: 5
      17 <212> TYPE: PRT
      18 <213> ORGANISM: Homo sapiens
     21 Thr Tyr Trp Leu Gly see ven 3 on Even Surmary Sheet
22 1
 --> 19 <400> SEQUENCE: 1
E--> 22 1
     56 <210> SEQ ID NO: 5
     57 <211> LENGTH: \7
     LENGTH: 15

235 <212> TYPE: DNA

236 <213> ORGANISM: Homo sapiens

237 <400> SEQUENCE: 10 | 5
      58 <212> TYPE: PRT
                                                The type of errors shown
W--> 60 <400> SEQUENCE: 5
E--> 63 1 <del>5</del>
W--> 237 <400> SEQUENCE: 10
                                          33 nucleo tides indicated
42 nucleo traces found
E--> 238 agatatacta tgcac
     255 <210> SEQ ID NO: 13
     256 <211> LENGTH: (38)
     257 <212> TYPE: DNA
     258 <213> ORGANISM: Homo sapiens
V--> 259 <400> SEQUENCE: 13
   260 ctctcctgca gggccagtca gagtgttagc agctacttag cc
                                                                                33
     262 <210> SEQ ID NO: 14
     263 <211> LENGTH: (21
     264 <212> TYPE: DNA
                                Errored: Bequence 14
27 nucleotides indicated
21 nucleotides found
     265 <213> ORGANISM: Homo sapiens
W--> 266 <400> SEQUENCE: 14
E--> 267 gatgcatcca acagggcc
                                                                                18
     269 <210> SEQ ID NO: 15
```

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/920,262

270 <211> LENGTH: 27

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/920,262

DATE: 08/09/2001

TIME: 14:31:25

Input Set : A:\PTO_VSK.txt

Output Set: N:\CRF3\08092001\I920262.raw

21 mulestides found

271 <212> TYPE: DNA

272 <213> ORGANISM: Homo sapiens

W--> 273 <400> SEQUENCE: 15

274 cagcagcgta gcaactggcc t E--> 278 CEN 248 4

E--> 280 1 E--> 283

21

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/920,262

DATE: 08/09/2001 TIME: 14:31:26

Input Set : A:\PTO_VSK.txt

Output Set: N:\CRF3\08092001\I920262.raw

L:11 M:270 C: Current Application Number differs, Replaced Current Application No L:11 M:271 C: Current Filing Date differs, Replaced Current Filing Date L:19 M:283 W: Missing Blank Line separator, <400> field identifier L:22 M:332 E: (32) Invalid/Missing Amino Acid Numbering, SEQ ID:1 L:30 M:283 W: Missing Blank Line separator, <400> field identifier L:42 M:283 W: Missing Blank Line separator, <400> field identifier L:51 M:283 W: Missing Blank Line separator, <400> field identifier L:60 M:283 W: Missing Blank Line separator, <400> field identifier L:63 M:332 E: (32) Invalid/Missing Amino Acid Numbering, SEQ ID:5 L:69 M:283 W: Missing Blank Line separator, <400> field identifier L:78 M:283 W: Missing Blank Line separator, <400> field identifier L:108 M:283 W: Missing Blank Line separator, <400> field identifier L:135 M:283 W: Missing Blank Line separator, <400> field identifier L:237 M:283 W: Missing Blank Line separator, <400> field identifier L:238 M:254 E: No. of Bases conflict, LENGTH:Input: 0 Counted:15 SEQ:10' L:245 M:283 W: Missing Blank Line separator, <400> field identifier L:252 M:283 W: Missing Blank Line separator, <400> field identifier L:259 M:283 W: Missing Blank Line separator, <400> field identifier L:260 M:254 E: No. of Bases conflict, LENGTH:Input:33 Counted:42 SEQ:13 L:260 M:252 E: No. of Seq. differs, <211>LENGTH:Input:33 Found:42 SEQ:13 L:266 M:283 W: Missing Blank Line separator, <400> field identifier L:267 M:252 E: No. of Seq. differs, <211>LENGTH:Input:21 Found:18 SEQ:14. L:273 M:283 W: Missing Blank Line separator, <400> field identifier L:278 M:340 E: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:15L:278 M:254 E: No. of Bases conflict, LENGTH:Input:4 Counted:24 SEQ:15 L:278 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1 L:278 M:112 C: (48) String data converted to lower case, M:254 Repeated in SeqNo=15 L:283 M:252 E: No. of Seq. differs, <211>LENGTH:Input:27 Found:24 SEQ:15